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1: Cancer Res. 199/ Aug 1;5/(15):3281-/.

Entrez PubMed

Cloning of P2XM, a novel human P2X receptor gene regulated by p53.

PubMed Services

Urano T, Nishimori H, Han H, Furuhata T, Kimura Y, Nakamura Y, Tokino T.

Laboratory of Molecular Medicine, The Institute of Medical Science, The University of Tokyo, Minato-ku, Japan.

Through cloning of functional p53-binding sites (p53-tagged sites) from the human genome, we isolated a novel gene inducible by wild-type p53. Its cDNA sequence contained an open reading frame encoding a 431-amino acid

peptide that showed a significant homology with members of the P2X family. This protein also revealed a similarity to RP-2, a gene activated in thymocytes undergoing programmed cell death. Northern blot analysis showed that it was expressed predominantly in skeletal muscle. Hence, we designated the gene P2XM (P2X specifically expressed in skeletal muscle). P2XM was localized to chromosomal band 22q11, where frequent loss of heterozygosity has been observed in rhabdoid tumors. Although we detected

a minor splice variant lacking a part of exon 1 that would encode residues corresponding to transmembrane domain M1 was relatively more abundant in two of seven sarcoma cell lines, one of which was derived from a rhabdomyosarcoma, and the other was derived from an osteosarcoma. The results suggest that P2XM may play a significant role in the proliferation and/or differentiation of skeletal muscle cells and that its altered expression may be involved in the development of some sarcomas.

no genetic alteration in the coding sequences, one of four rhabdomyosarcom? cell lines examined had completely lost expression of this gene. Furthermore

PMID: 9242461 [PubMed - indexed for MEDLINE]

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Related Resources

L4 ANSWER 11 OF 12 USPATFULL on STN

ACCESSION NUMBER: 2000:138518 USPATFULL TITLE: Purinergic receptor

INVENTOR(S): Buell, Gary Nutter, Geneva, Switzerland

Surprenant, Annmarie, Geneva, Switzerland

Kawashima, Eric, Geneva, Switzerland

PATENT ASSIGNEE(S): Glaxo Group Limited, Greenford, United Kingdom

(non-U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION:
APPLICATION INFO.:

US 6133434 US 1997-842079 20001017 19970428 (8)

DOCUMENT TYPE: FILE SEGMENT: Utility Granted

PRIMARY EXAMINER:

Spector, Lorraine

LEGAL REPRESENTATIVE:

Nixon & Vanderhye P.C.

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 10 1

NUMBER OF DRAWINGS:

47 Drawing Figure(s); 18 Drawing Page(s)

LINE COUNT:

1652

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

DETD The proteins, polypeptides and peptides of the invention can be used as antigens to generate P2X.sub.7 specific antibodies.

Methods of antibody generation are well known in the art. Both monoclonal and polyclonal antibodies are contemplated, as are antigen.

DETD . . . with PBS containing 5% bovine serum albumin, 5% goat serum with 1% Triton X-100. After blocking, sections were incubated with anti-P2X.sub.7 antibody at 10 mg/ml for 2 h at room temperature or 16 h at 4.degree. C., then washed in PBS. The. .

The specificity of the anti-P2X.sub.7
antibody was tested by Western blotting. Cells were harvested
with PBS containing 10 mM EDTA, washed twice and resuspended in PBS.
. under reducing conditions. Separated proteins were transferred to a
nitrocellulose membrane (Novex, San Diego, Calif.). The membrane was
incubated with anti-P2X.sub.7 antibody,
followed by incubation with peroxidase-coupled sheep anti-rabbit IgG
(Dako, Denmark) and developed using the ECL system (Amersham,
Buckinghamshire, UK). Proteins. . .

anti bidus

L11 ANSWER 20 OF 25 USPATFULL on STN

ACCESSION NUMBER: 2001:51805 USPATFULL

TITLE: Nucleic acids encoding a functional human

purinoreceptor P2X3 and P2X6, and methods of production

and use thereof

INVENTOR(S): Lynch, Kevin J., Gurnee, IL, United States

Burgard, Edward C., Libertyville, IL, United States

van Biesen, Tim, Chicago, IL, United States

PATENT ASSIGNEE(S): Abbott Laboratories, Abbott Park, IL, United States

(U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: APPLICATION INFO.:

US 6214581 B1 20010410 US 1998-191136 19981113 (9)

RELATED APPLN. INFO.:

US 1998-191136 19981113 (9) Continuation-in-part of Ser. No. US 1998-8526, filed on

16 Jan 1998, now abandoned Continuation-in-part of Ser. No. US 1998-8185, filed on 16 Jan 1998, now abandoned

NUMBER DATE

PRIORITY INFORMATION:

US 1998-71298P 19980116 (60)

US 1998-71669P

19980116 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT:

Granted Ulm, John

PRIMARY EXAMINER: LEGAL REPRESENTATIVE:

Becker, Cheryl L., Goller, Mimi C.

NUMBER OF CLAIMS:

14 1

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

16 Drawing Figure(s); 16 Drawing Page(s)

LINE COUNT:

2829

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Human P2X.sub.3 and P2X.sub.6 purinergic receptor

polypeptides are provided. Nucleic acid molecules encoding the aforementioned human P2X receptor polypeptide, and vectors and host

cells containing such.

DETD . . . above P2X receptors but also methods for screening compounds

using the receptor and cells expressing the receptor. Further, polynucleotides and antibodies which can be used in methods

for detection of the receptor, as well as the reagents useful in these

methods,.

DETD . . . of Ligand Binding, Wiley-Liss, Inc., N.Y.; Michel et al. (1997)

Mol. Pharmacol. 51:524-532). Alternatively, expression can be detected

by utilizing antibodies or functional measurements, i.e.,
ATP-stimulated cellular depolarization using methods that are well known

to those skilled in the art. For.

DETD Furthermore, each specific P2X polypeptide or fragment(s) thereof can be

used to prepare monoclonal antibodies using techniques that

are well known in the art. The specific P2X receptor or relevant fragments can be obtained using. . . specific P2X polypeptide or

fragments can be obtained using. . . specific F2x polypeptide of fragment(s) thereof can be synthesized using conventional polypeptide

synthetic techniques as known in the art. Monoclonal antibodies that display specificity and selectivity for a particular P2X

polypeptide can be labeled with a measurable and detectable moiety, for.

fluorescent moiety, radiolabels, enzymes, chemiluminescent labels and the like, and used in in vitro assays. It is theorized that such antibodies could be used to identify wild-type or variant P2X

receptor polypeptides for immuno-diagnostic purposes. For example, antibodies have been generated to detect amyloid b1-40 v. 1-42

in brain tissue (Wisniewski et al. (1996) Biochem. J. 313:575-580; also.

DETD . . . e.g., Winzor et al. (1995) Quantitative Characterization of Ligand Binding, Wiley-Liss, Inc., N.Y.). Alternatively, expression can

be detected by utilizing antibodies or functional measurements, i.e., ATP- or UTP-stimulated cellular depolarization using methods that are well known to those skilled in the. . .

L24 ANSWER 27 OF 33 USPATFULL on STN

ACCESSION NUMBER: 2001:29315 USPATFULL

TITLE: Method of screening for compounds that bind P2x

receptor

INVENTOR(S): Valera, Soledad, Geneva, Switzerland

Buell, Gary N, Geneva, Switzerland

PATENT ASSIGNEE(S): Glaxo Group Limited, Greenford, United Kingdom

(non-U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 6194162 B1 20010227

APPLICATION INFO.: US 1999-363745 19990730 (9)

RELATED APPLN. INFO.: Continuation of Ser. No. US 750134, now patented, Pat.

No. US 5985603

NUMBER DATE

PRIORITY INFORMATION: GB 1994-10664 19940527

GB 1995-2480 19950209

DOCUMENT TYPE: Utility

FILE SEGMENT: Granted
PRIMARY EXAMINER: Mertz, Prema

ASSISTANT EXAMINER: Murphy, Joseph F. LEGAL REPRESENTATIVE: Nixon & Vanderhye P.C.

NUMBER OF CLAIMS: 6
EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 29 Drawing Figure(s); 19 Drawing Page(s)

LINE COUNT: 1063

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 13 OF 33 PCTFULL COPYRIGHT 2003 Univentio on STN L24 1998018916 PCTFULL ED 20020514 ACCESSION NUMBER: TITLE (ENGLISH): HUMAN P2X PURINORECEPTOR TITLE (FRENCH): PURINO-RECEPTEUR P2X HUMAIN INVENTOR(S): HILLMAN, Jennifer, L.; COLEMAN, Roger INCYTE PHARMACEUTICALS, INC.; PATENT ASSIGNEE(S): HILLMAN, Jennifer, L.; COLEMAN, Roger English LANGUAGE OF PUBL.: DOCUMENT TYPE: Patent PATENT INFORMATION: DATE NUMBER KIND ______ WO 9818916 A1 19980507 DESIGNATED STATES AT AU BR CA CH CN DE DK ES FI GB IL JP KR MX NO NZ RU W: SE SG US GH KE LS MW SD SZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH DE DK ES FI FR GB GR IE IT LU MC NL PT SE BF BJ CF CG CI CM GA GN ML MR NE SN TD TG WO 1997-US18370 A 19971015 APPLICATION INFO.: US 1996-8/742,621 19961030 PRIORITY INFO.: HUMAN P2X PURINORECEPTOR TIEN PURINO-RECEPTEUR P2X HUMAIN TIFR The present invention provides a novel human P2X ABEN purinoreceptor (HPURR) and polynucleotides which identify and encode HPURR. The invention also provides genetically engineered expression vectors and host cells. ABFR Cette invention concerne un nouveau purino-recepteur P2X humain (HPURR), ainsi que des nucleotides permettant d'identifier et de coder ce HPURR. Cette invention concerne egalement des vecteurs d'expression. . HPURR in infected host cells DETD (Logan and Shenk (1984) Proc. Nad. Acad. Sci. 81:3655-3659). In addition, transcription enhancers, such as the Rous sarcoma virus (RSV) enhancer, may be used to increase expression in mammalian host cells. ion transport, signal transmission, and apoptosis. Such diseases include, but are not limited to, chronic pain, neuropathic pain such as diabetic-, cancer-, and AIDS-related, neurodegenerative diseases such as Alzheimer's disease, Parkinson's disease, Huntington's disease, Creutzfeld-Jacob disease, and amyotrophic lateral sclerosis, and dementias, including AIDS-related, as. al. (I 983) Immunol. Today 4:72; Cote et al. (1983) Proc. Natl. Acad. Sci. 80:2026-2030; Cole et al. (1985) Monoclonal Antibodies and Cancer Therap , Alan R. Liss Inc.,

For any compound, the therapeutically effective dose can be estimated initially either in cell culture assays, e.g., of **neoplastic** cells, or in animal models, usually mice, rabbits, dogs, or pigs. The animal model may also be used to determine the. . .

New York, NY, pp. 77-96).

.. ...,

osteoarthritis, asthma, systemic lupus, myasthenia gravis, diabetes mellitus, osteoporosis, glomerulonephritis, and scleroderma; neurological diseases including chronic pain, neuropathic pain such as diabetic-, cancer-, and AIDS-related, neurodegenerative diseases such as Alzheimer's disease, Parkinson's disease, Creutzfeld-Jacob disease, and amyotrophic lateral sclerosis, and dementias, such as AIDS-related,. . .

L24 ANSWER 26 OF 33 USPATFULL on STN

ACCESSION NUMBER: 2001:51805 USPATFULL

TITLE: Nucleic acids encoding a functional human

purinoreceptor P2X3 and P2X6, and methods of production and use thereof

INVENTOR(S): Lynch, Kevin J., Gurnee, IL, United States

Burgard, Edward C., Libertyville, IL, United States

van Biesen, Tim, Chicago, IL, United States

PATENT ASSIGNEE(S): Abbott Laboratories, Abbott Park, IL, United States

(U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 6214581 B1 20010410

APPLICATION INFO.: US 1998-191136 19981113 (9)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1998-8526, filed on 16 Jan 1998, now abandoned Continuation-in-part of Ser.

No. US 1998-8185, filed on 16 Jan 1998, now abandoned

NUMBER DATE

PRIORITY INFORMATION: US 1998-71298P 19980116 (60)

US 1998-71669P 19980116 (60)

DOCUMENT TYPE: Utility
FILE SEGMENT: Granted

PRIMARY EXAMINER: Ulm, John LEGAL REPRESENTATIVE: Becker, Cheryl L., Goller, Mimi C.

NUMBER OF CLAIMS: 14 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 16 Drawing Figure(s); 16 Drawing Page(s)

LINE COUNT: 2829

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 19 OF 25 USPATFULL on STN

ACCESSION NUMBER: 2001:82541 USPATFULL

TITLE: Nucleic acids encoding a functional human

purinoreceptor P2X2 and P2X4, and methods of production

and use thereof

INVENTOR(S): Lynch, Kevin J., Gurnee, IL, United States

Burgard, Edward C., Libertyville, IL, United States

Metzger, Randy E., Gurnee, IL, United States Niforatos, Wende, Chicago, IL, United States Touma, Edward B., Chicago, IL, United States Van Biesen, Tim, Chicago, IL, United States

PATENT ASSIGNEE(S): Abbott Laboratories, Abbott Park, IL, United States

(U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 6242216 B1 20010605

APPLICATION INFO.: US 1998-191608 19981113 (9)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1998-137458, filed

on 20 Aug 1998

NUMBER DATE

PRIORITY INFORMATION: US 1997-65822P 19971114 (60)

DOCUMENT TYPE: Utility
FILE SEGMENT: Granted
PRIMARY EXAMINER: Ulm, John

LEGAL REPRESENTATIVE: Becker, Cheryl L., Goller, Mimi C.

NUMBER OF CLAIMS: 20 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 29 Drawing Figure(s); 15 Drawing Page(s)

LINE COUNT: 1329

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Human P2X.sub.2 and P2X.sub.4 purinergic receptor polypeptides are provided. Nucleic acid molecules encoding the aforementioned human P2X receptor polypeptide, and vectors and host cells containing such. . .

DETD . . . above P2X receptors but also methods for screening compounds using the receptor and cells expressing the receptor. Further, polynucleotides and antibodies which can be used in methods for detection of the receptor, as well as the reagents useful in these methods, . . .

DETD Michel et al. (1997) Mol. Pharmacol. 51:524-532). Alternatively, expression can be detected by utilizing antibodies or functional measurements, i.e., ATP-stimulated cellular depolarization using methods that are well known to those skilled in the art. For.

Furthermore, each specific P2X polypeptide or fragment(s) thereof can be used to prepare monoclonal antibodies using techniques that are well known in the art. The specific P2X receptor or relevant fragments can be obtained using. . . specific P2X polypeptide or fragment(s) thereof can be synthesized using conventional polypeptide synthetic techniques as known in the art. Monoclonal antibodies that display specificity and selectivity for a particular P2X polypeptide can be labeled with a measurable and detectable moiety, for. . . fluorescent moiety, radiolabels, enzymes, chemiluminescent labels and the like, and used in in vitro assays. It is theorized that such antibodies could be used to identify wild-type or variant P2X receptor polypeptides for immuno-diagnostic purposes. For example, antibodies have been generated to detect amyloid b1-40 v. 1-42 in brain tissue (Wisniewski et al. (1996) Biochem. J. 313:575-580; also.

COPYRIGHT 2003 Univentio on STN ANSWER 10 OF 25 PCTFULL

1991016056 PCTFULL ED 20020513 ACCESSION NUMBER:

USE OF PURINERGIC RECEPTOR AGONISTS TITLE (ENGLISH):

AS ANTINEOPLASTIC AGENTS

EMPLOI D'AGONISTES DE RECEPTEUR PURINERGIQUE UTILISES TITLE (FRENCH):

COMME AGENTS ANTINEOPLASTIQUES

TREPEL, Jane, B.; INVENTOR(S): FANG, Wei-Gang;

PIRNIA, Farzaneh;

MYERS, Charles, E., Jr.

THE UNITED STATES OF AMERICA, as represented by THE PATENT ASSIGNEE(S):

SECRETARY, U.S. DEPARTMENT OF COMMERCE

LANGUAGE OF PUBL.: English

Patent DOCUMENT TYPE:

PATENT INFORMATION:

NUMBER KIND DATE ______ A1 19911031

WO 9116056 DESIGNATED STATES

> AT AU BE CA CH DE DK ES FR GB GR IT JP LU NL SE W:

APPLICATION INFO.: WO 1991-US1552 A 19910312

US 1990-509,183 19900416 PRIORITY INFO.:

TIEN USE OF PURINERGIC RECEPTOR AGONISTS AS

ANTINEOPLASTIC AGENTS

DETD . . agonists. For

example, specimen cells of such cancers can be tested for expression of P2 purinergic receptors by use of labeled ligand, anti-receptor antibody, or by a molecular probe, In addition, cells from fresh specimens can be assayed for Ca 2+ mobilization in response to P2.

19 8 9) . Cel 1 s were incubated at room temperature for 30-60 min.,, washed,, and resuspended in calcium medium. Rabbit anti-fluorescein antibody (1 unit/ml, Molecular Probes) was added to the cell suspension to reduce fluorescence background.

CLMEN. . . P2 purinergic

receptor agonists, comprising testing specimen cells of said cancer for expression of P. purinergic receptors by use of labeled ligand,, anti-receptor antibody, or a molecular probe.

17* A method for determining analogs or derivatives of purinergic receptor agonists therapeutically useful in treating hormone-independent cancers, comprising testing for. .

PCTFULL COPYRIGHT 2003 Univentio on STN ANSWER 10 OF 25 L11

1991016056 PCTFULL ED 20020513 ACCESSION NUMBER:

USE OF **PURINERGIC RECEPTOR** AGONISTS TITLE (ENGLISH):

AS ANTINEOPLASTIC AGENTS

EMPLOI D'AGONISTES DE RECEPTEUR PURINERGIQUE UTILISES TITLE (FRENCH):

COMME AGENTS ANTINEOPLASTIQUES

INVENTOR (S): TREPEL, Jane, B.; FANG, Wei-Gang;

PIRNIA, Farzaneh;

MYERS, Charles, E., Jr.

THE UNITED STATES OF AMERICA, as represented by THE PATENT ASSIGNEE(S):

SECRETARY, U.S. DEPARTMENT OF COMMERCE

LANGUAGE OF PUBL.: English

DOCUMENT TYPE: Patent

PATENT INFORMATION:

NUMBER KIND DATE ______

WO 9116056 A1 19911031

DESIGNATED STATES

AT AU BE CA CH DE DK ES FR GB GR IT JP LU NL SE W:

APPLICATION INFO.: WO 1991-US1552 A 19910312

PRIORITY INFO.: US 1990-509,183 19900416

PCTFULL COPYRIGHT 2003 Univentio on STN ANSWER 7 OF 12 L4

1995033048 PCTFULL ED 20020514 ACCESSION NUMBER:

P2X RECEPTORS (PURINOCEPTOR FAMILY) TITLE (ENGLISH):

RECEPTEURS P2X (FAMILLE DES PURINORECEPTEURS) TITLE (FRENCH):

INVENTOR (S): VALERA, Soledad; BUELL, Gary, Nutter

PATENT ASSIGNEE(S): GLAXO GROUP LIMITED;

VALERA, Soledad;

BUELL, Gary, Nutter

LANGUAGE OF PUBL.: English

DOCUMENT TYPE: Patent

PATENT INFORMATION:

NUMBER KIND DATE

______ WO 9533048 A2 19951207

DESIGNATED STATES

₩: AM AT AU BB BG BR BY CA CH CN CZ DE DK EE ES FI GB GE

HU IS JP KE KG KP KR KZ LK LR LT LU LV MD MG MN MW MX NO NZ PL PT RO RU SD SE SG SI SK TJ TM TT UA US UZ VN KE MW SD SZ UG AT BE CH DE DK ES FR GB GR IE IT LU MC

NL PT SE BF BJ CF CG CI CM GA GN ML MR NE SN TD TG A 19950524

WO 1995-EP1968 APPLICATION INFO.: PRIORITY INFO.:

GB 1994-9410664.8 19940527

GB 1995-9502480.8 19950209 L4 ANSWER 11 OF 12 USPATFULL on STN

ACCESSION NUMBER: 2000:138518 USPATFULL

TITLE: Purinergic receptor

INVENTOR(S): Buell, Gary Nutter, Geneva, Switzerland Surprenant, Annmarie, Geneva, Switzerland

Kawashima, Eric, Geneva, Switzerland

PATENT ASSIGNEE(S): Glaxo Group Limited, Greenford, United Kingdom

(non-U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 6133434 20001017

APPLICATION INFO.: US 1997-842079 19970428 (8)

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Spector, Lorraine

LEGAL REPRESENTATIVE: Nixon & Vanderhye P.C.

NUMBER OF CLAIMS: 10 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 47 Drawing Figure(s); 18 Drawing Page(s)

LINE COUNT: 1652

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

COPYRIGHT 2003 Univentio on STN ANSWER 8 OF 12 PCTFULL 1981000261 PCTFULL ED 20020506 ACCESSION NUMBER: CHARGE EFFECTS IN IMMUNOASSAYS TITLE (ENGLISH): EFFETS DE CHARGES DANS LES IMMUNOANALYSES TITLE (FRENCH): GIBBONS I; INVENTOR(S): ULLMAN E; ROWLEY G SYVA CO PATENT ASSIGNEE(S): English LANGUAGE OF PUBL.: DOCUMENT TYPE: Patent PATENT INFORMATION: NUMBER KIND DATE ______ WO 8100261 A1 19810205 DESIGNATED STATES W: DE GB JP A 19800423 APPLICATION INFO .: WO 1980-US455 PRIORITY INFO.: US 1979-61099 19790726 DETD (P2K2) 5or (P2X2) 5 Immunoglobulin D(IgD) or yD-Globulin (yD) VII Proconvertin VIII Antihemophilic globulin (AHG)

or yD-Globulin (yD)
VII Proconvertin
VIII Antihemophilic globulin
(AHG)
Ix Christmas factor,
plasma thromboplastin
component (PTC)
A U
OMPI
x Stuart-Prower factor,
autoprothrombin III
xi Plasma thromboplastin
antecedent (PTA)
xii Hagemann. factor
XIII Fibrin@stabilizing. . .

ANSWER 20 OF 86 BIOTECHDS COPYRIGHT 2003 THOMSON DERWENT/ISI on STN

ACCESSION NUMBER: 1998-11352 BIOTECHDS

TITLE:

Human gene P2XM whose transcription is induced by

p53;

useful for diagnostic purposes and in development of new

anticancer drugs

AUTHOR:

Tokino T; Nakamura Y

PATENT ASSIGNEE:

Otsuka-Pharm.

LOCATION:

Tokyo, Japan.

PATENT INFO:

WO 9842835 1 Oct 1998

APPLICATION INFO: WO 1998-JP1146 18 Mar 1998

PRIORITY INFO:

JP 1997-93044 26 Mar 1997

DOCUMENT TYPE: LANGUAGE:

Patent Japanese

OTHER SOURCE:

WPI: 1998-532006 [45]

development of new anticancer drugs. (43pp)

AN

1998-11352 BIOTECHDS

AB

A new human gene encoding a specified protein sequence which is significantly homologous to the P2X family of ATP receptors and the RP-2 protein which is expressed in T-lymphocytes during apoptosis is claimed. Transcription of the genes is specifically regulated by the tumor suppressor gene p53. Also claimed is a P2XM gene, which is specifically expressed in skeletal muscle and which has been localized to chromosome-22q11, an area where mutation and sequence losses frequently occur in rhabdoid sarcomas: The genes may be used for diagnostic purposes (e.g. by detecting changes in the gene in sarcomas), using DNA probes and DNA primers containing or derived

from all or part of the genes. The genes may further be used in the

PXM = PZX6

L34 ANSWER 30 OF 73 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN

ACCESSION NUMBER:

1999:466 BIOSIS PREV199900000466

DOCUMENT NUMBER: TITLE:

Expression of functional P2-purinergic receptors in primary

cultures of human colorectal carcinoma cells.

AUTHOR (S):

Hopfner, M.; Lemmer, K.; Jansen, A.; Hanski, C.; Riecken,

E.-O.; Gavish, M.; Mann, B.; Buhr, H.; Glassmeier, G.;

Scherubl, H.

CORPORATE SOURCE:

Abteilung Innere Medizin/Gastroenterol.,

Universitaetsklinikum Benjamin Franklin, Freie Univ.

Berlin, Hindenburgdamm 30, 12200 Berlin Germany

SOURCE:

Biochemical and Biophysical Research Communications, (

Oct. 29, 1998) Vol. 25, No. 3, pp. 811-817.

ISSN: 0006-291X.

DOCUMENT TYPE:

Article English

LANGUAGE:

L34 ANSWER 41 OF 73 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN

ACCESSION NUMBER:

CORPORATE SOURCE:

1999:112647 BIOSIS PREV199900112647

DOCUMENT NUMBER: TITLE:

Cytotoxic effects of ATP against B cell neoplasias

and acute myeloid leukaemia resulting from selective

expression of surface P2Z (P2x7) purinergic

receptors.

AUTHOR(S):

Lammas, D. A.; Quibell, K.; Kumararatne, D.; Drayson, M. Dep. Immunol., Med. Sch., Univ. Birmingham, Birmingham B15

2TT UK

SOURCE:

Immunology, (Dec., 1998) Vol. 95, No. SUPPL. 1,

pp. 31.

Meeting Info.: 6th Annual Congress of the British Society for Immunology Harrogate, England, UK December 1-4, 1998

ISSN: 0019-2805.

DOCUMENT TYPE:

Conference English

LANGUAGE:

L34 ANSWER 39 OF 73 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN

ACCESSION NUMBER: 1998:527623 BIOSIS

DOCUMENT NUMBER:

PREV199800527623

TITLE:

Development of purinergic receptors of P2x

-subtype on the megakaroyctic cells (Meg-01) derived from a

leukemia patient.

AUTHOR(S):

Kawa, Kazuyoshi (1)

CORPORATE SOURCE:

(1) Dep. Neurophysiol., Tohoku Univ. Sch. Med., 2-1

Seiryo-cho, Sendai, Miyagi 980-8575 Japan

SOURCE:

Neuroscience Research Supplement, (1998) No. 22, pp. S92.

Meeting Info.: 21st Annual Meeting of the Japan

Neuroscience Society and the First Joint Meeting of the Japan Neuroscience Society and the Japanese Society for Neurochemistry Tokyo, Japan September 21-23, 1998 Japan

Neuroscience Society . ISSN: 0921-8696.

DOCUMENT TYPE:

Conference English

LANGUAGE:

L34 ANSWER 37 OF 73 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STN

ACCESSION NUMBER: 1998:202482 SCISEARCH

THE GENUINE ARTICLE: ZA381

TITLE: Cytolytic P2X purinoceptors

AUTHOR: DiVirgilio F (Reprint); Chiozzi P; Falzoni S; Ferrari D;

Sanz J M; Venketaraman V; Baricordi O R

CORPORATE SOURCE: UNIV FERRARA, DEPT EXPT & DIAGNOST MED, SECT GEN PATHOL,

VIA BORSARI, 46, I-44100 FERRARA, ITALY (Reprint); UNIV FERRARA, CTR BIOTECHNOL, I-44100 FERRARA, ITALY; UNIV FERRARA, DEPT EXPT & DIAGNOST MED, MED GENET SECT, I-44100

FERRARA, ITALY

COUNTRY OF AUTHOR:

ITALY

SOURCE: CELL

CELL DEATH AND DIFFERENTIATION, (MAR 1998) Vol.

5, No. 3, pp. 191-199.

Publisher: STOCKTON PRESS, HOUNDMILLS, BASINGSTOKE,

HAMPSHIRE, ENGLAND RG21 6XS.

ISSN: 1350-9047.

DOCUMENT TYPE:

General Review; Journal

FILE SEGMENT:

LIFE

LANGUAGE:

English

REFERENCE COUNT:

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

L45 ANSWER 15 OF 86 MEDLINE on STN DUPLICATE 10

ACCESSION NUMBER: 1999366710 MEDLINE

DOCUMENT NUMBER: 99366710 PubMed ID: 10440098

TITLE: Pharmacological characterization of recombinant human and

rat P2X receptor subtypes.

AUTHOR: Bianchi B R; Lynch K J; Touma E; Niforatos W; Burgard E C;

Alexander K M; Park H S; Yu H; Metzger R; Kowaluk E; Jarvis

M F; van Biesen T

CORPORATE SOURCE: Neurological and Urological Diseases Research,

Pharmaceutical Products Division, Abbott Laboratories,

Abbott Park, IL 60064-3500, USA.

SOURCE: EUROPEAN JOURNAL OF PHARMACOLOGY, (1999 Jul 2)

376 (1-2) 127-38.

Journal code: 1254354. ISSN: 0014-2999.

PUB. COUNTRY: Netherlands

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199909

ENTRY DATE: Entered STN: 19991012

Last Updated on STN: 19991012 Entered Medline: 19990927







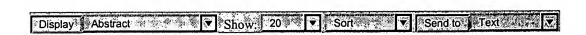
3	MCRI		uD ₁	Mea	of Medicine NLM					
Entrez	PubMed	Nucleotide	Protein	Genome	Structure	PMC	Journals	Во		
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<u></u>	— J	Display Abstract	<u> </u>	Show: 20	Sort	Send	to Text			
Entrez Pr	ubMed	1: Pharmacol FREE full te. www.pha	xt article at	Mar;53(1):107	7-18.	Rela	ated Articles,	Links		
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PubMed	Services		, Burnstock Iumphrey l	k G, Kennedy PP.	C, King BF	, North R	A, Seguela	P,		
				6-29, Californ mrc-lmb.cam		f Technol	ogy, Pasader	na,		
Related I	Resources	receptors the metabotrop	nat mediate of the properties	mediator to cothe actions of ptors and the	ATP belong transmitter-g	to two clarated, ion c	sses, the channel P2X			

ATP acts as a humoral mediator to control cell function extracellularly. The receptors that mediate the actions of ATP belong to two classes, the metabotropic P2Y receptors and the transmitter-gated, ion channel P2X receptors. This review describes the structure, distribution, function, and ligand recognition characteristics of P2X receptors, which comprise seven distinct subunits that can function as both homo- and hetero- polymers. The pharmacology of P2X receptors is complicated by marked differences between species orthologues. The current nomenclature is based largely on recombinant receptor studies and detailed knowledge of endogenous P2X receptors in native tissues is limited because of lack of good selective agonists and antagonists for each receptor type.

Publication Types:

- Review
- Review, Academic

PMID: 11171941 [PubMed - indexed for MEDLINE]



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Department of Health & Human Services
Freedom of Information Act | Disclaimer

L24 ANSWER 26 OF 33 USPATFULL on STN

ACCESSION NUMBER: 2001:51805 USPATFULL

Nucleic acids encoding a functional human TITLE:

> purinoreceptor P2X3 and P2X6, and methods of production and use thereof Lynch, Kevin J., Gurnee, IL, United States

INVENTOR (S):

Burgard, Edward C., Libertyville, IL, United States

van Biesen, Tim, Chicago, IL, United States

Abbott Laboratories, Abbott Park, IL, United States PATENT ASSIGNEE(S):

(U.S. corporation)

NUMBER KIND DATE ______

US 6214581 B1 20010410 PATENT INFORMATION: US 1998-191136 19981113 (9) APPLICATION INFO.:

Continuation-in-part of Ser. No. US 1998-8526, filed on RELATED APPLN. INFO.: 16 Jan 1998, now abandoned Continuation-in-part of Ser.

No. US 1998-8185, filed on 16 Jan 1998, now abandoned .

DATE NUMBER

US 1998-71298P 19980116 (60) PRIORITY INFORMATION:

19980116 (60) US 1998-71669P

DOCUMENT TYPE: Utility FILE SEGMENT: Granted Ulm, John PRIMARY EXAMINER:

Becker, Cheryl L., Goller, Mimi C. LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: 14 EXEMPLARY CLAIM: 1

16 Drawing Figure(s); 16 Drawing Page(s) NUMBER OF DRAWINGS:

LINE COUNT: 2829

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Nucleic acids encoding a functional human purinoreceptor P2X3 and P2X6, and methods of production and use thereof

Human P2X.sub.3 and P2X.sub.6 purinergic receptor AB polypeptides are provided. Nucleic acid molecules encoding the aforementioned human P2X receptor polypeptide, and vectors and host cells containing such nucleic acid molecules, are also provided. In addition, methods are provided for producing these P2X

receptor polypeptide, as are methods of using such polypeptides and host cells that express the same to screen for compounds having activity on P2X.sub.3 and P2X.sub.6 receptors. Further,

therapeutic uses involving aspects of these receptors are contemplated.

. supra, pp 337-345), immune and inflammatory diseases (Di SUMM Virgilio et al. (1995) in: Belardinelli et al. (eds), supra, pp 329-335), cancer (Rapaport (1993) Drug Dev. Res. 28:428-431), constipation and diarrhea (Milner et al. (1994) in: Kamm et al. (eds.) Constipation and.

. supra, pp 337-345, immune and inflammatory diseases (Di SUMM Virgilio et al. (1995) in: Belardinelli et al. (eds), supra, pp 329-335), cancer (Rapaport (1993) Drug Dev. Res. 28:428-431), constipation and diarrhea (Milner et al. (1994) in: Kamm et al. (eds.) Constipation and.

. . of a polynucleotide into a prokaryotic cell. "Transformation" DETD of a eukaryotic cell also may refer to the formation of a cancerous or tumorigenic state.

. cells also are known in the art and include viral promoters DETD such as that from Simian Virus 40 (SV40), Rous sarcoma virus (RSV), adenovirus (ADV), bovine papilloma virus (BPV) and cytomegalovirus (CMV). Mammalian cells also may require terminator sequences and poly.

DETD . . . identical to the human P2X6 receptor described herein which is expressed at high levels in skeletal muscle (Urano et al. Cancer

Res. 57:3281-3287 (1997)). Additionally, this gene is inducible by the p53 tumor suppressor gene product, suggesting that the human P2X.sub.6 receptor plays a role in skeletal muscle cell proliferation and/or differentiation. Therefore, agents that modulate the activity of the P2X.sub.6 receptor may be useful as therapeutics for musculoskeletal disorders such as sarcomas. The practice of the present invention will employ, unless otherwise indicated, conventional techniques of molecular biology, microbiology, recombinant DNA technology, . . .

DETD

. . . of a polynucleotide into a prokaryotic cell. "Transformation" of a eukaryotic cell also may refer to the formation of a cancerous or tumorigenic state.

DETD

. . . cells also are known in the art and include viral promoters such as that from Simian Virus 40 (SV40), Rous **sarcoma** virus (RSV), adenovirus (ADV), bovine papilloma virus (BPV) and cytomegalovirus (CMV). Mammalian cells also may require terminator sequences and poly. . .

PubMed Entrez BLAST OMIM Taxonomy Structure

BLAST 2 SEQUENCES RESULTS VERSION BLASTP 2.2.6 [Apr-09-2003]

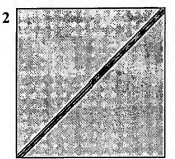
Matrix BLOSUM62 ▼ gap open: 11 gap extension: 1

x_dropoff: 50 expect: 10.0 wordsize: 3 Filter Align

Sequence 1 lcl|seq_1 Length 431 (1 .. 431)

Sequence 2 lcl|seq_2 **Length** 431 (1 .. 431)





PZXM PZX6

NOTE: The statistics (bitscore and expect value) is calculated based on the size of nr database

Score = 913 bits (2360), Expect = 0.0 Identities = 431/431 (100%), Positives = 431/431 (100%)

MGSPGATTGWGLLDYKTEKYVMTRNWRVGALQRLLQFGIVVYVVGWALLAKKGYQERDLE 60 Query: 1 MGSPGATTGWGLLDYKTEKYVMTRNWRVGALQRLLQFGIVVYVVGWALLAKKGYQERDLE MGSPGATTGWGLLDYKTEKYVMTRNWRVGALQRLLQFGIVVYVVGWALLAKKGYQERDLE 60 Sbjct: 1 POFSIITKLKGVSVTOIKELGNRLWDVADFVKPPQGENVFFLVTNFLVTPAQVQGRCPEH 120 Query: 61 PQFSIITKLKGVSVTQIKELGNRLWDVADFVKPPQGENVFFLVTNFLVTPAQVQGRCPEH Sbjct: 61 PQFSIITKLKGVSVTQIKELGNRLWDVADFVKPPQGENVFFLVTNFLVTPAQVQGRCPEH 120 Query: 121 PSVPLANCWVDEDCPEGEGGTHSHGVKTGQCVVFNGTHRTCEIWSWCPVESGVVPSRPLL 180 PSVPLANCWVDEDCPEGEGGTHSHGVKTGQCVVFNGTHRTCEIWSWCPVESGVVPSRPLL Sbjct: 121 PSVPLANCWVDEDCPEGEGGTHSHGVKTGQCVVFNGTHRTCEIWSWCPVESGVVPSRPLL 180 Query: 181 AQAQNFTLFIKNTVTFSKFNFSKSNALETWDPTYFKHCRYEPQFSPYCPVFRIGDLVAKA 240 AQAQNFTLF1KNTVTFSKFNFSKSNALETWDPTYFKHCRYEPQFSPYCPVFR1GDLVAKA Sbjct: 181 AQAQNFTLFIKNTVTFSKFNFSKSNALETWDPTYFKHCRYEPQFSPYCPVFRIGDLVAKA 240 Query: 241 GGTFEDLALLGGSVGIRVHWDCDLDTGDSGCWPHYSFQLQEKSYNFRTATHWWEQPGVEA 300 GGTFEDLALLGGSVGIRVHWDCDLDTGDSGCWPHYSFQLQEKSYNFRTATHWWEQPGVEA Sbjct: 241 GGTFEDLALLGGSVGIRVHWDCDLDTGDSGCWPHYSFQLQEKSYNFRTATHWWEQPGVEA 300 Query: 301 RTLLKLYGIRFDILVTGQAGKFGLIPTAVTLGTGAAWLGVVTFFCDLLLLYVDREAHFYW 360 RTLLKLYGIRFDILVTGQAGKFGLIPTAVTLGTGAAWLGVVTFFCDLLLLYVDREAHFYW Sbjct: 301 RTLLKLYGIRFDILVTGQAGKFGLIPTAVTLGTGAAWLGVVTFFCDLLLLYVDREAHFYW 360 Query: 361 RTKYEEAKAPKATANSVWRELALASQARLAECLRRSSAPAPTATAAGSQTQTPGWPCPSS 420 RTKYEEAKAPKATANSVWRELALASQARLAECLRRSSAPAPTATAAGSQTQTPGWPCPSS Sbjct: 361 RTKYEEAKAPKATANSVWRELALASQARLAECLRRSSAPAPTATAAGSQTQTPGWPCPSS 420

0.02 total secs.

Query: 421 DTHLPTHSGSL 431 DTHLPTHSGSL Sbjct: 421 DTHLPTHSGSL 431 0.00 sys. secs CPU time: 0.02 user secs. Lambda K 0.137 0.452 0.321 Gapped Lambda K 0.267 0.0410 0.140 Matrix: BLOSUM62 Gap Penalties: Existence: 11, Extension: 1 Number of Hits to DB: 2003 Number of Sequences: 0 Number of extensions: 131 Number of successful extensions: 1 Number of sequences better than 10.0: 1 Number of HSP's better than 10.0 without gapping: 1 Number of HSP's successfully gapped in prelim test: 0 Number of HSP's that attempted gapping in prelim test: 0 Number of HSP's gapped (non-prelim): 1 length of query: 431 length of database: 498,883,957 effective HSP length: 130 effective length of query: 301 effective length of database: 498,883,827 effective search space: 150164031927 effective search space used: 150164031927 T: 9 A: 40 X1: 16 (7.4 bits) X2: 129 (49.7 bits) X3: 129 (49.7 bits) S1: 41 (21.8 bits) S2: 76 (33.9 bits)

L24 ANSWER 16 OF 33 PCTFULL COPYRIGHT 2003 Univentio on STN ACCESSION NUMBER: 1995033048 PCTFULL ED 20020514

TITLE (ENGLISH): 1995033048 FC1F011 ED 20020514

P2X RECEPTORS (PURINOCEPTOR FAMILY)

TITLE (FRENCH): RECEPTEURS P2X (FAMILLE DES PURINORECEPTEURS)

INVENTOR(S): VALERA, Soledad; BUELL, Gary, Nutter PATENT ASSIGNEE(S): GLAXO GROUP LIMITED;

> VALERA, Soledad; BUELL, Gary, Nutter

LANGUAGE OF PUBL.: English

DOCUMENT TYPE: Patent

PATENT INFORMATION:

NUMBER KIND DATE
----WO 9533048 A2 19951207

DESIGNATED STATES

TaT .

AM AT AU BB BG BR BY CA CH CN CZ DE DK EE ES FI GB GE HU IS JP KE KG KP KR KZ LK LR LT LU LV MD MG MN MW MX NO NZ PL PT RO RU SD SE SG SI SK TJ TM TT UA US UZ VN KE MW SD SZ UG AT BE CH DE DK ES FR GB GR IE IT LU MC NL PT SE BF BJ CF CG CI CM GA GN ML MR NE SN TD TG

APPLICATION INFO.: WO 1995-EP1968
PRIORITY INFO.: GB 1994-9410664

WO 1995-EP1968 A 19950524 GB 1994-9410664.8 19940527 GB 1995-9502480.8 19950209

TIEN P2X RECEPTORS (PURINOCEPTOR FAMILY)

TIFR RECEPTEURS P2X (FAMILLE DES PURINORECEPTEURS)

ABEN The P2X receptor of ATP has been cloned and expressed by recombinant DNA technology, so the receptor can be prepared free from other ATP receptors. The P2X receptor enables antibodies to be prepared and is useful in screening compounds for use in a variety of diseases and. . .

ABFR Le recepteur P2X d'adenosine 5'-triphosphate (ATP) a ete clone et exprime par technique de recombinaison d'ADN, de sorte que le recepteur puisse etre obtenu separement, sans les autres recepteurs d'ATP. Le recepteur P2X permet de preparer des anticorps, et peut etre utilise pour detecter des composes a utiliser dans une variete de maladies. .

DETD . . . of hP 2X from

urinary bladder isolation of human P2X cDNA Human urinary bladder tissue was obtained from a cystectomy for a bladder tumor. The patient showed no symptoms of bladder instability or urodynamic abnormalities. Only those portions, surrounding the tumor, which appeared macroscopically normal (Palea et al - supra) were used. Total RNA was isolated by guanidinium isothiocyanate and poly A' RNA.

analysis (Figure 7). HL60 cells can be differentiated into distinct lineages, depending on the inductant (Koeffler, Induction of Differentiation of Human Acute Myelogenous Leukemia Cells: Therapeutic Implications Blood 62: 709-721 (1983)). Induction of macrophage-like characteristics with phorbol diesters or granulocytic differentiation with DMSO or dibutryl.

L56 ANSWER 14 OF 17 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 1992:81141 CAPLUS

DOCUMENT NUMBER: 116:81141

TITLE: P2-purinergic receptor agonists inhibit the growth of

androgen-independent prostate carcinoma cells Fang, Wei Gang; Pirnia, Farzaneh; Bang, Yung Jue;

AUTHOR(S): Fang, Wei Gang; Pirnia, Farzaneh; Bang, Yung
Myers, Charles E.; Trepel, Jane B.

CORPORATE SOURCE: Div. Cancer Treat., Natl. Cancer Inst., Bethesda, MD,

20892, USA

SOURCE: Journal of Clinical Investigation (1992),

89(1), 191-6

CODEN: JCINAO; ISSN: 0021-9738

DOCUMENT TYPE: Journal LANGUAGE: English

To develop a new approach to the treatment of advanced, hormone-refractory prostate cancer, the signal transduction regulating the growth of human androgen-independent prostate carcinoma cell lines were studied. Agonist-stimulated Ca2+ mobilization, a crit. regulatory event in other secretory cell types, was studied as a means of identifying previously undescribed plasma membrane receptors that may transduce a growth inhibitory signal. In all of the cell lines tested, P2-purinergic receptor agonists, including ATP and certain hydrolysis-resistant adenine nucleotides, induced a rapid, transient increase in cytoplasmic free Ca2+ that was detectable at 50 to 100 nM ATP, was maximal at 100 .mu.M ATP, and was inhibited .apprx.50% by chelation of extracellular Ca2+. Within 8 s after addn., ATP stimulated accumulation of the polyphosphatidylinositol products inositol 1,4,5-trisphosphate, inositol 1,3,4-trisphosphate, and inositol tetrakisphosphate. In addn. to stimulating phosphatidylinositol turnover and Ca2+ mobilization, ATP and hydrolysis-resistant ATP analogs induced >90% inhibition of the growth of all lines tested. These data demonstrate that human androgen-independent prostate carcinoma cells express functional P2-purinergic receptors linked to phospholipase C, and that agonists of this receptor are markedly growth inhibitory, suggesting a novel therapeutic approach to this common adult neoplasm.

L56 ANSWER 15 OF 17 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 1991:623457 CAPLUS

DOCUMENT NUMBER: 115:223457

TITLE: Use of purinergic receptor agonists as antineoplastic

agents

INVENTOR(S): Trepel, Jane B.; Fang, Wei-Gang; Pirnia, Farzaneh;

Myers, Charles E., Jr.

PATENT ASSIGNEE(S): National Institutes of Health, USA

SOURCE: U. S. Pat. Appl., 33 pp. Avail. NTIS Order No.

PAT-APPL-6-509 183.

CODEN: XAXXAV

DOCUMENT TYPE:

LANGUAGE:

F.

Patent English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 509183	A0	19910301	US 1990-509183	19900416 <
WO 9116056	A1	19911031	WO 1991-US1552	19910312 <
W: AU, CA,	JP			
RW: AT, BE,	CH, DE	, DK, ES,	FR, GB, GR, IT, LU, NL	, SE
AU 9175792	A1	19911111	AU 1991-75792	19910312 <
US 5415873	A	19950516	US 1993-131033	19931004 <
US 5641500	Α	19970624	US 1995-446954	19950515 <
PRIORITY APPLN. INFO	.:		US 1990-509183	19900416
			WO 1991-US1552	19910312
			US 1992-888292	19920526
			US 1993-131033	19931004

Purinergic receptor agonists inhibit hormone-independent, including AB androgen-independent, cancer cells which express the purinergic receptor. Methods to screen for advanced cancers that are amenable to such treatment and for new, potent analogs or derivs. of purinergic receptor agonists are also disclosed. A wide variety of surface receptors that induced an increase in Ca2+ was found in human prostate carcinoma cell lines. ATP produced the largest Ca2+ responses of the agonists tested. This response was so large that there was almost complete depletion of internal Ca2+ stores. Cell lines DU145, PC-3, and PC-3M expressed a P2 subtype purinergic receptor with agonist potency ATP > ADP > AMP > adenosine. Nonhydrolyzable analogs .alpha.,.beta.-methylene ATP, .beta.,.gamma.-methylene ATP, 5'-adenylylimidodiphosphate (AMP-PNP), and adenosine -5'-0-(3thiophosphate) had agonist activity. ATP and AMP-PNP inhibited growth of the above 3 cell lines. The cell growth was assocd. with an increase in DNA strand breaks.

L62 ANSWER 21 OF 22 USPATFULL on STN

ACCESSION NUMBER: 95:43035 USPATFULL

TITLE: Use of purinergic receptor agonists as antineoplastic

agents

INVENTOR(S): Trepel, Jane B., Bethesda, MD, United States

Fang, Wei-Gang, Bethesda, MD, United States Pirnia, Farzaneh, Potomac, MD, United States

Myers, Jr., Charles E., Rockville, MD, United States

PATENT ASSIGNEE(S): The United States of America as represented by the

Department of Health and Human Services, Washington,

DC, United States (U.S. government)

NUMBER KIND DATE

PATENT INFORMATION: US 5415873 19950516

APPLICATION INFO.: US 1993-131033 19931004 (8)

RELATED APPLN. INFO.: Continuation of Ser. No. US 1992-888292, filed on 26

May 1992, now abandoned which is a continuation of Ser. No. US 1990-509183, filed on 16 Apr 1990, now abandoned

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Kishore, Gollamudi S. LEGAL REPRESENTATIVE: Morgan & Finnegan

NUMBER OF CLAIMS: 11 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 18 Drawing Figure(s); 13 Drawing Page(s)

LINE COUNT: 603

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

SUMM The present invention relates to a method for treating hormone-independent cancers via the use of purinergic receptor agonists, diagnostic uses of these compounds to determine effective treatment for specific tumors, a

process for screening for new potent analogs of these compounds, and the

use of these compounds in facilitating the.

SUMM

. . . an androgen-dependent cell line. By incubating the cells in vitro with an agonist of a known cell surface receptor and detecting an intracellular response, it became possible to screen for a wide variety of receptors previously unreported on these cells. Ca.sup.2+. . . and Ca.sup.2+ regulated channels. Interestingly, there have not been any reported studies of this signal transduction system in benign or neoplastic prostatic tissue. Thus, the results disclosed here are the first report of (1) hormone-stimulated Ca.sup.2+ transients in prostatic cells; (2) hormone-stimulated phosphatidylinositol turnover in prostatic cells; (3) purinergic receptor expression on prostatic cells; and

(4) purinergic receptor-associated cell death in

prostatic adenocarcinoma.

L45 ANSWER 9 OF 86 MEDLINE on STN DUPLICATE 5

ACCESSION NUMBER: 2001547250 MEDLINE

DOCUMENT NUMBER: 21477952 PubMed ID: 11593539

TITLE: P2Z purinoceptor, a special receptor for

apoptosis induced by ATP in human leukemic

lymphocytes.

AUTHOR: Peng L; Bradley C J; Wiley J S

CORPORATE SOURCE: Department of Laboratory Medicine, First Hospital, West

China University of Medical Sciences, Chengdu 610041,

China.

SOURCE: CHINESE MEDICAL JOURNAL, (1999 Apr) 112 (4)

356-62.

Journal code: 7513795. ISSN: 0366-6999.

PUB. COUNTRY:

China
Journal; Article; (JOURNAL ARTICLE)

DOCUMENT TYPE: LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

200112

ENTRY DATE: Entered STN: 20011015

Last Updated on STN: 20020122 Entered Medline: 20011207

OBJECTIVE: To investigate the role of purinergic P2Z receptors AB for apoptosis of human leukemic lymphocytes mediated by extracellular adenosine triphosphate (ATP). METHODS: A total of 13 B-chronic lymphocytic leukemia (CLL) patients were studied. Exposure of leukemic lymphocytes with (n = 8) or without (n = 5)P2Z receptors to ATP, benzoylbenzoic-ATP (BzATP), 2-methylthio-ATP (2MeSATP), adenosine-5' [gamma-thio] triphosphate (ATP-gamma S), and other nucleosides for 8 h in vitro. Apoptosis was detected by electron microscopy (EM), agarose gel electrophoresis, and the quantitative assay-TdT assay. RESULTS: Apoptosis was detected only in leukemic lymphocytes with P2Z receptors. Using a quantitative assay, ATP-induced DNA strand breaks were found to occur specifically with BzATP, ATP and 2MeSATP, but not for analogue ATP-gamma S nor other nucleosides. Meanwhile, ATP-induced DNA fragmentation was fully blocked by pretreatment with oxidized ATP (OxATP), a compound recently shown to block P2Z receptors. Also, it is shown that the Ca2+/calmodulin complex plays a role in the regulation of the apoptosis induced by ATP on CLL cells, because an antagonist of this complex, 1-[N, O-bis (5isoquinolinesulfonyl)-N-methyl-L-tyrosyl]-4-phenylpiperazine (KN-62) was found to inhibit the ATP-induced apoptosis. Furthermore, choline, an inhibitor of phospholipase D (PLD), is first shown to partially inhibit ATP-induced apoptosis. CONCLUSION: These data indicate that P2Z receptors on lymphocytes play an important role in the apoptosis induced by ATP in vitro.

L45 ANSWER 11 OF 86 MEDLINE on STN DUPLICATE 7

ACCESSION NUMBER: 1999145440 MEDLINE

DOCUMENT NUMBER: 99145440 PubMed ID: 9989927

TITLE: Activation of the P2Z/P2X7 receptor in

human lymphocytes produces a delayed permeability lesion:

involvement of phospholipase D.

AUTHOR: Fernando K C; Gargett C E; Wiley J S

CORPORATE SOURCE: Sydney University Department of Medicine, The Nepean

Hospital, Somerset Street, Penrith, Australia.

SOURCE: ARCHIVES OF BIOCHEMISTRY AND BIOPHYSICS, (1999 Feb

15) 362 (2) 197-202.

Journal code: 0372430. ISSN: 0003-9861.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199903

ENTRY DATE: Entered STN: 19990324

Copyright 1999 Academic Press.

Last Updated on STN: 19990324 Entered Medline: 19990311

AB Leukemic lymphocytes possess a cytolytic P2Z/

P2X7 receptor which, when activated by extracellular ATP, opens a Ca2+- and Ba2+-permeable ion channel. This ATP-stimulated influx of divalent cations has been shown to activate an intracellular phospholipase D (PLD) which hydrolyzes membrane phosphatidylcholine. Lymphocytes that were exposed to ATP for 20 min at 37 degrees C, washed, and then incubated without ATP for 2 h showed an increased uptake of propidium2+, a dye widely used to measure cytotoxicity. The potent P2Z/ P2X7 receptor inhibitor, KN-62, which is known to prevent the channel opening when added with ATP, did not block development of the permeability lesion when added 15 min before dye addition. The activity of lymphocyte PLD was stimulated fourfold by ATP and a proportion of this increased activity persisted for several hours after removal of ATP. Loading lymphocytes with intracellular choline+ by prior incubation of cells with ATP in isotonic choline chloride abolished both ATP-stimulated PLD activity and the ATP-induced permeability lesion. Addition of PLD but not phospholipase C to the extracellular medium increased lymphocyte permeability to propidium2+ and this effect was not observed in a choline medium. The cytolytic effect of exogenous PLD together with the inhibitory effect of choline, a product of the PLD reaction, suggests that sustained activation of intracellular PLD may be involved in the ATP-initiated cytolytic pathway.

L45 ANSWER 20 OF 86 BIOTECHDS COPYRIGHT 2003 THOMSON DERWENT/ISI on STN

ACCESSION NUMBER: 1998-11352 BIOTECHDS

TITLE: Human gene P2XM whose transcription is induced by

p53;

useful for diagnostic purposes and in development of new

APPLICATION NO. DATE

anticancer drugs

AUTHOR: Tokino T; Nakamura Y

PATENT ASSIGNEE: Otsuka-Pharm. LOCATION: Tokyo, Japan.

PATENT INFO: WO 9842835 1 Oct 1998

APPLICATION INFO: WO 1998-JP1146 18 Mar 1998 PRIORITY INFO: JP 1997-93044 26 Mar 1997

DOCUMENT TYPE: Patent LANGUAGE: Japanese

OTHER SOURCE: WPI: 1998-532006 [45]

L45 ANSWER 21 OF 86 CAPLUS COPYRIGHT 2003 ACS on STN

ACCESSION NUMBER: 1998:650172 CAPLUS

DOCUMENT NUMBER: 129:326959

TITLE: Cloning of cDNA of a novel p53-inducible human gene

P2XM

INVENTOR(S): Tokino, Takashi; Nakamura, Yusuke

PATENT ASSIGNEE(S): Otsuka Pharmaceutical Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 16 pp.

KIND DATE

CODEN: JKXXAF

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.

		-					-				
JР	10262681		A2	19981006		JP 1997-93044		19970326	<		
WO	9842835		A1	19981001		WO 1998-JP1146	5	19980318	<		
	W: AU,	CA,	CN, ID,	KR, MX,	SG,	US					
	RW: AT,	BE,	CH, DE,	DK, ES,	FI,	FR, GB, GR, IE,	IT,	LU, MC,	ΝL,	PT,	SE
AU	9864184		A1	19981020		AU 1998-64184		19980318	<		
AU	724681		B2	20000928		•					
EP	1006186					EP 1998-909733					
	R: AT,	BE,	CH, DE,	DK, ES,	FR,	GB, GR, IT, LI,	LU,	, NL, SE,	MC,	PT,	
	IE,	FI									
US	6255472		B1	20010703		US 2000-381681		20000110			
PRIORIT	Y APPLN.	INFO	. :		Ú	JP 1997-93044	Α	19970326			
					Ţ	NO 1998-JP1146	W	19980318			

L24 ANSWER 15 OF 33 PCTFULL COPYRIGHT 2003 Univentio on STN

ACCESSION NUMBER: 1997023218 PCTFULL ED 20020514

TITLE (ENGLISH): A PROCESS FOR REGULATING VAGAL TONE TITLE (FRENCH): PROCEDE POUR REGULER LE TONUS VAGAL

INVENTOR(S): PELLEG, Amir PATENT ASSIGNEE(S): PELLEG, Amir

LANGUAGE OF PUBL.: English
DOCUMENT TYPE: Patent

PATENT INFORMATION:

DESIGNATED STATES

W: AU CA JP AT BE CH DE DK ES FI FR GB GR IE IT LU MC NL

PT SE

APPLICATION INFO.: WO 1996-US20255 A 19961224 PRIORITY INFO.: US 1995-60/009,228 19951226 US 1996-8/771,581 19961223

ABEN . . . invention provides methods of altering vagal tone in a patient by administering a

therapeutically effective amount of a mediator of P2x -purinoceptors located on vagal afferent nerve

terminals to the patient. Diagnostic applications are also provided.

ABFR . . tonus vagal chez un patient, en lui administrant une quantite suffisante, pour avoir un effet therapeutique, d'un mediateur des

purinorecepteurs P2x situes sur les terminaisons afferentes du nerf vague. L'invention concerne

egalement des utilisations diagnostiques.

DETD Patent No. 4,673,563. ATP has also been shown to be effective against cancer in animal models and in humans. However, the mechanism of action of ATP in this setting involves the immune system and/or direct action on tumor cells and is independent of the autonomic nervous system. The use of ATP as anti-cancer therapy is the subject of Rapaport, U.S. Patent No. 5,049,372.

ANSWER 2 OF 5 L65 ACCESSION NUMBER:

PCTFULL COPYRIGHT 2003 Univentio on STN

1998019541 PCTFULL ED 20020514

TITLE (ENGLISH):

METHODS AND COMPOSITIONS FOR TREATING AND DIAGNOSING

TUMORS USING ADENOSINE RECEPTOR ACTIVATED CELLS

TITLE (FRENCH):

PROCEDES ET COMPOSITIONS DE TRAITEMENT ET DE DIAGNOSTIC DE TUMEURS METTANT EN OEUVRE DES CELLULES ACTIVEES DU

RECEPTEUR DE L'ADENOSINE

INVENTOR(S):

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NEELY, Constance, F.

LANGUAGE OF PUBL.:

English

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Patent

PATENT INFORMATION:

PATENT ASSIGNEE(S):

NUMBER KIND DATE ______

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AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GE HU ID IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK TJ TM TR TT UA UG US UZ VN YU ZW GH KE LS MW SD SZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH DE DK ES FI FR GB GR IE IT LU MC NL PT SE BF BJ CF CG CI CM GA

GN ML MR NE SN TD TG

APPLICATION INFO.: PRIORITY INFO.:

WO 1997-US19926

19971105 Α

US 1996-8/748,559

19961108

Fang et al. reported the inhibition of cell growth in hormone-DETD refractory prostate cancer cell lines using P2 purinergic receptor agonists.

> ftinctionaIP2-purinergic receptors, and proposed that agonists of such receptors be used to inhibit the growth of related neoplasms. Methods of

treating prostate cancers by administration of a P2 purinergic receptor agonist

are provided in U.S. Patent 5,415,873.

L24 ANSWER 25 OF 33 USPATFULL on STN

ACCESSION NUMBER: 2001:82541 USPATFULL

TITLE: Nucleic acids encoding a functional human

purinoreceptor P2X2 and P2X4, and methods of production and use thereof Lynch, Kevin J., Gurnee, IL, United States

INVENTOR(S): Lynch, Kevin J., Gurnee, IL, United States

Burgard, Edward C., Libertyville, IL, United States

Metzger, Randy E., Gurnee, IL, United States Niforatos, Wende, Chicago, IL, United States Touma, Edward B., Chicago, IL, United States Van Biesen, Tim, Chicago, IL, United States

PATENT ASSIGNEE(S): Abbott Laboratories, Abbott Park, IL, United States

(U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 6242216 B1 20010605 APPLICATION INFO.: US 1998-191608 19981113 (9)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1998-137458, filed

on 20 Aug 1998

NUMBER DATE

PRIORITY INFORMATION: US 1997-65822P 19971114 (60)

DOCUMENT TYPE: Utility
FILE SEGMENT: Granted
PRIMARY EXAMINER: Ulm, John

LEGAL REPRESENTATIVE: Becker, Cheryl L., Goller, Mimi C.

NUMBER OF CLAIMS: 20 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 29 Drawing Figure(s); 15 Drawing Page(s)

LINE COUNT: 1329

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

TI Nucleic acids encoding a functional human purinoreceptor P2X2

and P2X4, and methods of production and use thereof Human P2X.sub.2 and P2X.sub.4 purinergic receptor

Human P2X.sub.2 and P2X.sub.4 purinergic receptor polypeptides are provided. Nucleic acid molecules encoding the aforementioned human P2X receptor polypeptide, and vectors and host cells containing such nucleic acid molecules, are also provided. In addition, methods are provided for producing these P2X receptor polypeptide, as are methods of using such polypeptides and host cells that express the same to screen for compounds having activity on P2X.sub.2 and P2X.sub.4 receptors. Further,

therapeutic uses involving aspects of these receptors are contemplated.

SUMM . . . supra, pp 337-345), immune and inflammatory diseases (Di Virgilio et al. (1995) in: Belardinelli et al. (eds), supra, pp 329-335), cancer (Rapaport (1993) Drug Dev. Res. 28:428-431), constipation and diarrhea (Milner et al. (1994) in: Kamm et al. (eds.) Constipation and . . .

DETD . . . of a polynucleotide into a prokaryotic cell. "Transformation" of a eukaryotic cell also may refer to the formation of a cancerous or tumorigenic state.

DETD . . . cells also are known in the art and include viral promoters such as that from Simian Virus 40 (SV40), Rous sarcoma virus (RSV), adenovirus (ADV), bovine papilloma virus (BPV) and cytomegalovirus (CMV). Mammalian cells also may require terminator sequences and poly. . .

DETD . . . identical to the human P2X.sub.6 receptor described herein which is expressed at high levels in skeletal muscle (Urano et al. Cancer Res. 57:3281-3287 (1997)). Additionally, this gene is inducible by the p53 tumor suppressor gene product, suggesting that the human P2X.sub.6 receptor plays a role in skeletal muscle cell proliferation and/or differentiation. Therefore, agents that modulate

the activity of this receptor may be useful as therapeutics for musculoskeletal disorders such as **sarcomas**.

L20 ANSWER 5 OF 8 USPATFULL on STN

ACCESSION NUMBER: 2001:51805 USPATFULL

TITLE: Nucleic acids encoding a functional human

purinoreceptor P2X3 and P2X6, and methods of production and use thereof

INVENTOR(S): Lynch, Kevin J., Gurnee, IL, United States

Burgard, Edward C., Libertyville, IL, United States

van Biesen, Tim, Chicago, IL, United States

PATENT ASSIGNEE(S): Abbott Laboratories, Abbott Park, IL, United States

(U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 6214581 B1 20010410

APPLICATION INFO.: US 1998-191136 19981113 (9)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1998-8526, filed on 16 Jan 1998, now abandoned Continuation-in-part of Ser.

No. US 1998-8185, filed on 16 Jan 1998, now abandoned

05 1996-6165, liled on 16 dan 1996, now abandone

NUMBER DATE

PRIORITY INFORMATION: US 1998-71298P 19980116 (60)

US 1998-71669P 19980116 (60)

DOCUMENT TYPE: Utility
FILE SEGMENT: Granted
PRIMARY EXAMINER: Ulm, John

LEGAL REPRESENTATIVE: Becker, Cheryl L., Goller, Mimi C.

NUMBER OF CLAIMS: 14
EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 16 Drawing Figure(s); 16 Drawing Page(s)

LINE COUNT: 2829

CAS INDEXING IS AVAILABLE FOR THIS PATENT.